

PCT/GB2004/004928

## 18 MAY 2006

1	"Detection of Protein Interactions"
2	
3	Field of the Invention
4	
5	The present invention relates to a method of
6	detecting interactions. In particular, but not
7	exclusively, the invention relates to a method of
8	detecting protein to protein interactions using
9	fluorescence.
10	·
11	Background to the Invention
12	
13	Protein to protein interactions play a key role in
14	many biological processes including the assembly of
15	enzymes, protein homo/hetero-oligomers, regulation
16	of intracellular transport, gene expression,
17	receptor-ligand interactions, entry of pathogens
18	into the cell and the action of small molecules or
19	drugs.
20	

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Identification and characterisation of 1 macromolecular interactions can be performed using 2 co-immunoprecipitation from cell lysates and 3 solubilised membranes. However, this technique 4 requires specific antibodies for both capture and 5 identification of proteins and may further require 6 the use of detergent to disrupt interactions. 7 8 More recently non-invasive techniques have been 9 developed to determine protein to protein 10 interactions. 11 12 Such non-invasive techniques were pioneered by the 13 yeast two hybrid method which is based on 14 complementation of a split yeast nuclear 15 transcription factor. 16 17 The use of yeast expression systems to identify 1.8 mammalian protein-to-protein interaction suffers 19 from a number of disadvantages. Certain post-20 translational modifications, that are normally 21 critical to mammalian protein interactions, cannot 22 be achieved by expression and / or post 23 translational modification of proteins by yeast 24 cells. For example, tyrosine phosphorylation is the 25 key to many mammalian intracellular protein binding 26 events involved in signal transduction. However, the 27 yeast genome contains no tyrosine kinase genes so 28 phosphotyrosine-dependent protein interactions 29 cannot be accessed in yeast two hybrid studies. 30 31

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1 Furthermore, in yeast two hybrid screening the 2 protein complex must be able to translocate to the 3 nucleus to cause expression of the reporter gene or 4 cause downstream events to trigger the expression of a reporter gene. Thus, proteins that are excluded 5 from the yeast nucleus will not be accessible to 6 this screening method. 7 8 Further methods such as protein complementation and 9 the split ubiquitin method utilise similar 10 11 underlying concepts to the yeast two hybrid method in that the interaction of two proteins (a bait 12 protein and prey protein) act to express a reporter 13 14 gene, the reporter gene allowing the interaction event to be visualised as a detectable signal. 15 16 Such methods which utilise the expression of a 17 reporter gene such as an enzyme to produce a 18 detectable signal suffer from the disadvantage that 19 the location of the protein complexes being detected 20 21 cannot be accurately visualised in the cell. 22 Recently the technique of fluorescence energy 23 24 transfer (FRET) has been used to determine protein to protein interactions. In this technique the 25 interaction of two fluorophores, an absorbing moiety 26 27 and a fluoresceing moiety, indicates their close spatial proximity. For protein to protein 28 interaction monitoring, the absorbing moiety is 29 30 added to a first protein partner and the fluorescing moiety is added to a second binding partner. 31 32 Provided the emission spectrum of the absorbing

moiety overlaps the excitation spectrum of the

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2 fluorescing moiety and both moieties are within 100Å 3 of each other FRET will occur. 4 5 FRET can utilise mutations in the sequence of green fluorescent protein (GFP) from the jellyfish 6 Aequorea victoria which have been shown to cause 7 variations in the spectral emission of GFP. 8 mutations give rise to variants of GFP such as 9 Yellow Fluorescent Protein (YFP), as well as cyan 10 (CFP) and blue (BFP) fluorescing variants. This 11 . technique uses fluorescent energy transfer between 12 these colour variants of GFP fused to interacting 13 proteins. Unfortunately, this method requires 14 overexpression of the GFP fusion proteins to allow 15 16 quantification of the small changes in fluorescence. Related methods to FRET require the use of 17 irreversible photobleaching (FRAP) or expensive 18 19 instruments capable of measuring fluorescence 20 lifetime imaging (FLIM). 21 22 It has recently been shown that fluorescence can be generated following the functional association of 23 24 two separate fragments of the GFP molecule (hapto-GFPs) when driven by the interaction of a pair of 25 proteins fused via a linker to the new C' and N' 26 termini of the hapto-GFPs. (Ghosh et al, (2000); Hu 27 et al, (2002). 28 29 30 Whilst the methods disclosed by these documents may be used in determining whether interaction occurs 31 32 between specific proteins they are not suitable for

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1 screening the interactions of peptides of which the mode of binding is unknown. 2 3 4 Conventionally, the length of the linkers used is chosen from a knowledge the peptides whose 5 6 interaction with each other is being tested. 7 this knowledge a suitable linker length which allows 8 association of the fragments of fluorescent protein 9 following the peptide interaction can be chosen. 10 knowledge of the peptides of interest or their mode of binding to each other has been considered to be 11 12 required. 13 14 For example, if the peptides interact with each other such that they form an anti-parallel complex 15 (hapto-GFP- $N^1$ -> $C^1$ :binding to : $C^2$ -> $N^2$ -hapto-GFP) and 16 the fluorescent fragments are orientated such that 17 18 they are directed away from each other in space then long linkers would be required to allow the 19 fragments of fluorescent protein to interact. 20 21 short linkers were used, despite interaction of the peptides of interest occurring, then this might not 22 be detected as the fragments would be prevented from 23 24 associating with each other due to the 25 stereochemical hindrance from the linkers. 26 would result in a false negative result in an assay 27 method. 28 29 30 31

1	Summary of the Invention
2	
3	The inventors through extensive work have developed
4	a robust system which overcomes many of the problems
5	of the prior art and provides for the first time a
6	general screening method which may used to determine
7	interaction between unknown peptides.
8	
9	According to a first aspect of the invention there
10	is provided a protein interaction system comprising
11	
12	a plurality of bait fusion proteins, each
13	fusion protein comprising (i) a first fragment
14	of fluorescent protein, a first peptide of
15	interest and a linker portion interposed
16	between the first peptide and first fluorescent
17	fragment; wherein the linker portions of each
18	bait fusion protein are of different lengths,
19	and the first peptide of interest of each bait
20	fusion protein is identical to the first
21	peptide of interest in each of the other bait
22	fusion proteins,
23	
24	and (ii) at least one prey fusion protein
25	comprising a fragment of fluorescent protein
26	complementary to said first fragment of
27	fluorescent protein, a second peptide of
28	interest and a second linker portion interposed
29	between the complementary fragment and the
30	second peptide;
31	

1	wherein, on interaction of a first peptide of
2	interest with a second peptide of interest, the
3	fragments of the fluorescent protein
4	functionally associate to promote fluorescence.
5	·
6	Thus, fluorescence will only be promoted when
7	peptides of interest of bait and prey fusion
8	proteins, having suitable linker lengths to allow
9	the respective fluorescent protein fragments to
10	associate, are used.
11:	
12	The provision of a peptide of interest linked to a
13	fluorescent fragment via a range of linker lengths
14	is advantageous over a single linker length as such
15	a range maximises the chances of an interaction
16	between peptides of interest being detected and
17	minimises the chances that the fluorescent fragments
18	cannot associate with each other due to
19	stereochemical hindrance or that the linkers are too
20	flexible (too long) and thus the fragments are not
21	being brought together in space despite the proteins
22	of interest interacting.
23	
24	The provision of fusion proteins wherein the fusion
25	proteins comprise linkers of different lengths
26	allows for the first time the provision of a general
27	method which can be used to study the interaction of
28	peptides of known and / or unknown structure and
29	also with bulkier peptides of interest and small
30	peptides of interest which interact with each other
31	such that the fragments of fluorescent protein are

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1 directed away from each other or peptides of unknown 2 structure. 3 4 Preferably at least three different linker lengths 5 are provided. More preferably at least four, even 6 more preferably at least five different linker 7 lengths are provided. 8 9 In an embodiment of the protein interaction system, 10 the system may additionally comprise at least one bait fusion protein which is identical to one of the 11 12 bait fusion proteins provided by the plurality of 13 bait fusion proteins. 14 15 A plurality of prey fusion proteins may be provided. 16 The linker portions of at least two prey fusion 17 proteins may be of different lengths. For example 18 two prey fusion proteins may be provided each 19 comprising the same protein of interest and same 20 fluorescent fragment, but provided with linkers of different lengths e.g. 10 amino acid residues and 20 21 22 amino acids respectively. 23 24 In one embodiment the linker portions comprise in 25 the range 5 to 60 amino acid residues, more 26 preferably in the range 5 to 60 amino acid, yet more preferably in the range 20 to 60 amino acid 27 residues. 28 29 30 In a preferred embodiment at least one of the linker 31 portions has at least 20 amino acids.

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1 In particular embodiments of the invention a linker 2 may comprise greater than 25 amino acids, for 3 example greater than 30 amino acids, greater than 35 4 amino acids, greater than 40 amino acids, greater 5 than 50 amino acids or greater than 55 amino acids 6 in length. 7 8 Preferably, the linker comprises up to 60 amino acids. 9 10 11 More preferably the linker comprises up to 45 amino 12 acids. 13 14 Preferably the linker is comprised of substantially hydrophillic amino-acid residues. 15 16 17 More preferably at least one, preferably each of the linkers comprises multiples of a pentapeptide 18 19 sequence such as glycyl-glycyl-glycyl-serine 20 (SEQ ID NO: 1). 21 22 Any fluorescent protein in which appropriate split sites can be formed and which the resulting 23 24 fragments can associate with each other and cause fluorescence may be used in the invention. Examples 25 26 of fluorescent proteins include red fluorescent protein and blue, yellow and cyan variants of GFP. 27 Moreover, variants of GFP which have increased 28 fluorescence may be utilised. However, in a 29 preferred embodiment the fragments of fluorescent 30 31 protein are fragments of green fluorescent protein, 32 mutants or variants thereof.

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2	More preferably the fluorescent protein is a	
3	humanised form of a fluorescent protein, e.g.	
4	Enhanced Green Fluorescent Protein (EGFP) or a	
5 .	variant thereof.	
6		
7	In a humanised nucleotide sequence one or more of	
8	the codons in the sequence are altered such that for	
9	the amino acid being encoded, the codon used is that	
LO	which most frequently appears in humans. This is	
11	advantageous as a humanised fluorescent protein	
12	construct e.g. (EGFP) has maximised expression	
13	levels and rate of flurophore formation in mammalian	
14	cells. This makes detection of fluorescence,	
15	produced by fragments of fluorescent proteins	
16	(fluorogenic fragments) which functionally associate	
17	with each other, easier to determine.	
18		
19	In preferred embodiments, the fragments of	
20	fluorescent protein (fluorogenic fragments) are	
21	generatable through the introduction of a split	
22	point between the amino acids at positions 157 and	
23	158, or (in a second embodiment) between the amino	
24	acids at positions 172 and 173 of the humanised form	
25	of Green Fluorescent Protein (SEQ ID NO 2) shown	
26	below.	
27		
28	SEQ ID NO 2 - EGFP (Clontech Inc.) [Genebank	
29	Accession number gb:AAB02574 gi 1377912]:	
30	1 mvskgeelft gvvpilveld gdvnghkfsv sgegegdaty	
31	41 gkltlkfict tgklpvpwpt lvttltygvq cfsrypdhmk	
32	81 qhdffksamp egyvqertif fkddgnyktr aevkfegdtl	

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121 vnrielkgid fkedgnilgh kleynynshn vyimadkqkn 1 161 gikvnfkirh niedgsvqla dhyqqntpig dgpvllpdnh 2 3 201 ylstqsalsk dpnekrdhmv llefvtaagi tlgmdelyk The fluorogenic fragments generated by the 5 6 introduction of a split point between the amino acid residues at positions 157 and 158, or between amino 7 acid residues at positions 172 and 173, result in 8 the production of hapto-EGFP1/157 and hapto-EGFP158/239, 9 or hapto-EGFP<sup>1/172</sup> and hapto-EGFP<sup>173/239</sup>, respectively. 10 11 Alternative split points are between residues 23/24, 12 38/39, 50/51, 76/77, 89/90, 102/103, 116/117, 13 132/133, 142/143, 190/191, 211/212 or 214/215 of 14 15 EGFP. 16 Thus in preferred embodiments, the fluorogenic 17 fragments are of amino acid residues 1 to 23, 1 to 18 38, 1 to 50, 1 to 76, 1 to 89, 1 to 102, 1 to 116, 1 19 to 132, 1 to 142, 1 to 157, 1 to 172, 1 to 190, 1 to 20 211 or 1 to 214, and a respective complementary 21 fragment 24 to 239, 39 to 239, 51 to 239, 77 to 239, 22 23 90 to 239, 103 to 239, 117 to 239, 133 to 239, 143 to 239, 158 to 239, 173 to 239, 191 to 239, 212 to 24 239, or 215 to 239 of EGFP. 25 26 It can be envisaged that three or more fluorescent 27 28 fragments may be provided by introducing two split 29 points as discussed above into the fluorescent protein, each fragment being fused to a peptide of 30 interest. On interaction of the peptides, the three 31 or more fluorescent fragments are brought together 32

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1 such that they can functionally associate and 2 generate a fluorescent signal capable of being 3 detected. 4 5 In another embodiment one or more of the three fluorescent fragments can be fused to a test agent 6 such as a small molecule, such as a metal ion. 7 this manner, protein interactions which require the 8 9 participation of additional test agents, such as small molecules, can be detected. 10 11 In an embodiment of the system wherein a plurality 12 13 of prey fusion proteins are present, additionally or 14 alternatively to prey proteins which comprise 15 linkers of different lengths at least two of the second peptides of interest of the prey fusion 16 17 proteins may comprise different amino acid 18 sequences. 19 20 The prey fusion peptides may be provided as a 21 library of different peptides of interest linked to 22 a fragment of fluorescent protein which is 23 complementary to the fluorescent fragment of the bait fusion protein. The library may be an 24 25 expression library, a library of a range of 26 mutations of a single peptide or other peptide 27 libraries as known in the art. 28 29 The first peptide of interest may be linked to the N terminus of the first fragment of fluorescent 30 31 protein or alternatively the first peptide may be

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linked to the C terminus of the first fragment of 1 2 fluorescent protein. 3 The second peptide of interest may be linked to the N terminus of the complementary fragment of 5 fluorescent protein or alternatively the second 6 7 peptide may be linked to the C terminus of the complementary fragment of fluorescent protein. 8 9 The peptides of interest linked to the fragments of 10 fluorescent protein can be small peptides of 11 12 differing amino acid sequence, for example nonomers, 13 comprising different amino acid compositions or the same overall composition, but with the amino acids 14 15 present in a different order. Alternatively, the peptides may be full size proteins e.g. obtained 16 from a cDNA library. Peptides may be produced 17 18 synthetically or recombinantly using techniques which are widely available in the art. For peptides 19 20 translated in the cell, naturally or induced, post-21 translational modification for example 22 glycosylation, lipidation, phosphorylation of the 23 peptides may occur, and these post translated products are still to be regarded as peptides. 24 25 26 In one embodiment, the protein interaction system is 27 a cell based interaction system. 28 29 In such a cell based system, each cell preferably 30 comprises one bait fusion protein of a single defined linker length. For example, if three bait 31 32 fusion proteins are provided each of which has a

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1 different linker length then a first cell will comprise a bait fusion protein of a first linker 2 3 length, a second cell will comprise a bait fusion 4 protein of a second linker length and a third cell will comprise a third bait fusion protein of a third 5 6 linker length. 7 When the protein interaction system is provided as a 8 9 cell based system, it may be produced using nucleic acid constructs which when expressed in live cells 10 provide the components of the protein interaction 11 12 system. 13 According to a second aspect of the present 14 invention there is provided a library of nucleic 15 acid constructs, each construct encoding 16 (i) a first fragment of fluorescent protein 17 18 capable of functional association with a complementary fragment of fluorescent protein 19 such that on functional association of said first 20 21 and complementary fragments fluorescence is 22 enabled, 23 (ii) a peptide of interest, and (iii) a linker portion interposed between the 24 peptide and first fragment of fluorescent protein 25 wherein the peptide of interest encoded by each 26 nucleic acid construct is the same and the linker 27 28 portion of each construct is of a different 29 length to the linker of each other construct. 30 31 In preferred embodiments at least one linker portion comprises at least 20 amino acids. 32

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2	The inventors have determined an economical and		
3	relatively easy way of providing longer (for example		
4	greater than 20 amino acids) linkers in the bait and		
5	/ or prey fusion proteins by providing linkers		
6	comprising multiples of a pentapeptide sequence such		
7	as glycyl-glycyl-glycyl-serine. Such		
8	sequences provide advantageous flexibility		
9	properties and thus enable the linker region to be		
<b>i</b> 0	readily extended to provide a robust screening		
11	method.		
12			
13	According to a third aspect of the invention there		
14	is provided an expression vector comprising a		
15	plurality of the constructs as provided by the		
16	second aspect of the invention.		
17			
18	According to a fourth aspect of the invention there		
19	is provided an expression vector comprising at least		
20	one of the plurality of nucleic acid constructs		
21	wherein the at least one nucleic acid construct		
22	encodes a fusion protein having a linker of at least		
23	20 amino acids.		
24			
25	An expression vector may be introduced into a cell		
26	using any known techniques such as calcium phosphate		
27	precipitation, lipofection, electroporation and the		
28	like.		
29			
30	In embodiments of the invention more than one vector		
31	encoding a construct of the third or fourth aspect		
32	of the invention and / or a construct comprising a		

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7	complementary fragment of fluorescent protein may be
2	introduced to a cell based system.
3	
4	According to a fifth aspect of the present invention
5	there is provided an assay method for monitoring
6	peptide interaction comprising the steps of
7	
8	providing the protein interaction system as
9	provided in the first aspect of the invention,
10	and
11	·
12	detecting fluorescence produced by the
13	interaction of first and second peptides of
14	interest causing fragments of the fluorescent
15	protein to functionally associate with each
16	other.
17	
18	In a particular embodiment the assay method is
19	performed in vitro.
20	
21	By providing fusion proteins of the protein
22	interaction system in a cell based system or by
23	mixing the fusion proteins of the first and second
24	protein of interest together in vitro the assay can
25	be used to screen a protein fusion library to
26	identify a second peptide of interest which binds to
27	a first peptide of interest or vice versa.
28	
29	An embodiment of the assay may comprise the step of
30	observing the subcellular location of the
31	interaction of the first and second peptides of
32	interest in a cell. This step is advantageous as it

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provides details of the location in the cell that 1 the interaction is taking place, for example at the 2 3 membrane, in the cytoplasm, or in the nucleus. 4 Any methods as known in the art may be used to 5 determine the subcellular location of interaction, 6 for example confocal scanning laser microscopy. 7 8 The assay method may further comprise the step of 9 observing the level of fluorescence produced at a 10 11 range of time points. 12 This step would allow determination of the 13 subcellular dynamics of the peptide interactions 14 visualised by fluorescence observations of living 15 16 cells to enable spatio-temporal studies of peptide interactions throughout all parts of the cell cycle, 17 for example such techniques would also allow the 18 trafficking of interacting peptides, for example 19 from the endoplasmic reticulum (ER) to the plasma 20 membrane to be tracked. 21 22 The assay may comprise the step of determining the 23 24 length of the linkers of those fusion proteins which allow the first fragment and complementary fragment 25 of the fluorescent protein to functionally 26 27 complement each other and enable fluorescence to be detected on interaction of the first and second 28 proteins of interest. 29 30 In such an embodiment the assay method may comprise 31 32 the steps of

**18** .

1 .	•
2	providing the protein interaction system as
3	provided in the first aspect of the invention,
4	
5	detecting fluorescence produced by the
6	interaction of the first and second peptides of
7	interest causing fragments of the fluorescent
8	protein to functionally associate with each
9	other,
10	
11	selecting those cells in which fluorescence is
12	detected,
13	
14	clonally amplifying the cells in which
15	fluorescence is detected, and
16	
17	determining the length of the linkers in said
18	cells by DNA sequencing.
19	·
20	Determination of the linker length of those fusion
21	proteins which interact with each other may be
22	advantageous as the distribution of occurrence of
23	linker lengths obtained from those cells in which
24	fluorescence is observed should indicate a sharp
25	cutoff at the lower limit of linker lengths
26	reflecting the minimum linker length capable of
27	spanning the separation of the fusion termini of the
28	interacting peptides. This in turn can be used to
29	provide a value of the distance between the
30	interacting peptides in Angstroms on the basis that
31	each amino acid residue contributes 3.7Å to the

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1 length of each linker in an extended backbone 2 . conformation. 3 4 An embodiment of the assay may comprise the further 5 step of isolating those fusion proteins which are 6 determined as allowing the first fragment and 7 complementary fragment of the fluorescent protein to 8 functionally complement each other and enable 9 fluorescence to be detected on interaction of the 10 first and second peptides of interest. 11 12 Isolation may be achieved for example using a 13 fluorescence activated cell sorting machine or laser 14 microdissection. 15 16 In a particular embodiment of this assay laser 17 excision of cell, amplification of the construct and 18 sequencing may be used to allow the linker lengths of those bait and / or prey fusion proteins of 19 20 interest which interact to cause fluorescence to be 21 determined and thus indicate the minimum distance of 22 the attachment points of the peptides of interest. 23 24 The isolated cells and fusion proteins may be 25 subjected to further analysis, for example 26 sequencing of the interacting peptides. 27 sequenced peptides may then be compared with 28 sequences (full length or partial) in a databank so as to identify or characterise the interacting 29 30 peptide isolated from the cell. 31

1	The sequences of the interacting peptides may	
2	alternatively be inferred by cloning selected	
3	fluorescent cells and subjecting the cloned cells to	
4	e.g. PCR amplification and DNA sequencing. These	
5	sequences can then be cloned into expression vectors	
6	and the protein expressed and purified. The	
7	purified protein can be further studied or used for	
8	example in research.	
9		
10	The assay may be used to determine if test agents	
11	are capable of promoting or enhancing interaction of	
12	peptides or alternatively of preventing or	
13	inhibiting the interaction of peptides.	
14		
15	In such an embodiment the assay may comprise the	
16	steps of	
17		
18	providing the protein interaction system as	
19	provided in the first aspect of the invention,	
20		
21	detecting the level of fluorescence produced by	
22	the interaction of the first and second	
23	peptides of interest causing fragments of the	
24	fluorescent protein to functionally complement	
25	each other,	
26		
27	providing a putative interaction modulating	
28	agent, and	
29		
30	detecting the level of fluorescence produced in	
31	the presence of said putative modulating agent,	
32		

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wherein detection of fluorescence in the 1 2 absence of the putative modulating agent, but not in the presence of the putative modulating 3 agent is indicative that the putative 4 modulating agent prevents or is an inhibitor of 5 peptide interaction and the detection of 6 fluorescence in the presence of the putative 7 modulating agent, but not in the absence of the 8 putative modulating agent is indicative that 9 the putative modulating agent promotes or 10 enhances peptide interaction. 11 12 The detected fluorescence may be quantitatively 13 determined such that fluorescence produced by 14 different cells or under different conditions can be 15 16 compared. 17 For example, in testing the effects of a putative 18 modulating agent, any detected fluorescence may be 19 20 measured in the absence and presence of the putative modulating agent wherein a reduction in fluorescence 21 22 in the presence of said modulating agent compared to fluorescence in the absence of said candidate 23 modulating agent is indicative of inhibition of 24 25 complex formation by the modulating agent and an increase in fluorescence is indicative of promotion 26 or enhancement of complex formation by the 27 28 modulating agent. 29 Modulation of the interaction between peptides may 30 31 be a desirable outcome in the treatment of certain clinical conditions, or as a research tool to study 32

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22 peptide to peptide interactions. For example, 1 modulation of peptide to peptide interactions may 2 facilitate the task of determining the steps of 3 complex pathways by the provision of means to 4 promote or inhibit a specific interaction, allowing 5 the effects of other proteins to be studied in 6 7 better detail. 8 9 Many peptide to peptide interactions require the 10 participation of small molecules or peptides. a requirement can be determined by simply adding 11 small molecules or peptides to a cell based system 12 or to an in vitro mixture containing the fusion 13 proteins of the interaction system and performing an 14 assay as described above to determine if these small 15 molecules or peptides modulate the interaction of 16 17 the peptides of interest as determined by detection or measurement of an alteration in fluorescent 18 19 signal. 20 21 Thus, embodiments of the assay of the present invention can be used to select compounds capable of 22 triggering, stabilising or destablising peptide to 23 24 peptide interactions. Embodiments of the assay method as described herein may be used to screen for 25 example, a receptor agonist, a receptor antagonist, 26 27 protein inhibitors, or an inhibitor of protein to protein interactions. 28 29 As will be apparent, the assay of the present 30

invention can be applied in a format appropriate for

large scale screening, for example, combinatorial

technologies can be employed to construct

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combinatorial libraries of small molecules or 2 peptides to test as modulating agents. 3 4 Preferably, structural information on the peptide to 5 peptide interaction to be modulated is obtained by 6 testing different agents to determine if they are 7 8 modulating agents. 9 For example, each of the interacting pair can be 10 expressed and purified and then allowed to interact 11 in suitable in vitro conditions. Optionally the 12 interacting peptides can be stabilised by 13 crosslinking or other techniques. The interacting 14 complex can be studied using various biophysical 15 techniques such as X-ray crystallography, NMR, or 16 mass spectrometry. In addition, information 17 concerning the interaction can be derived through 18 mutagenesis experiments for example alanine 19 scanning, or altering the charged amino acids or 20 hydrophobic residues on the exposed surface of the 21 bait or prey peptide being tested. 22 23 Based on the structural information obtained, 24 structural relationships between the interacting 25 peptides as well as between the modulating compound 26 and the interacting peptides can be elucidated. 27 Further, the three dimensional structure of the 28 interacting moieties and / or that of the modulating 29 compound can provide information to determine 30 suitable lead compounds able to modulate 31 interaction, which medicinal chemists can use to 32

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1 design analog compounds having similar moieties and 2 structures. 3 4 In a sixth aspect of the present invention there is provided novel compounds obtained using an assay of 5 6 the invention. 7 8 Modulator compounds obtained according to the method of invention may be prepared as a pharmaceutical 9 preparation or composition. 10 1:1 Such preparations will comprise the modulating 12 compound and a suitable carrier, diluent or 13 14 excipient. These preparations may be administered 1.5 by a variety of routes, for example, oral, buccal, 16 topical, intramuscular, intravenous, subcutaneous or 17 the like. 18 According to an seventh aspect of the present 19 invention there is provided a kit comprising nucleic 20 21 acid constructs as provided in the second aspect of the invention and means to express the constructs. 22 23 24 The kit may further comprise candidate modulating 25 agents, which promote, enhance, prevent or inhibit 26 peptide interaction. 27 The kit may further comprise nucleic acids which 28 encode at least one complementary fragment of 29 30 fluorescent protein, at least one second peptide of

interest and a second linker portion interposed

25

1 between the complementary fragment and the second 2 peptide of interest. 3 4 In another embodiment the kit comprises a cell in 5 which a vector comprising constructs of the second aspect of the invention can be expressed. 6 7 8 The kit may comprise a plurality of second peptides of interest of different amino acid sequences linked 9 to a complementary fragment of fluorescent protein. 10 11 12 Additionally, the kit may include instructions for 13 using the kit to practice the present invention. 14 The instructions should be in writing in a tangible 15 form or stored in an electronically retrievable 16 form. 17 18 Preferred features of each aspect of the invention 19 are as for each of the other aspects mutatis 20 mutandis unless the context demands otherwise. 21 22 Unless otherwise defined, all technical and 23 scientific terms used herein have the meaning 24 commonly understood by a person who is skilled in the art in the field of the present invention. 25 26 27 Throughout the specification, unless the context 28 demands otherwise, the terms 'comprise' or 29 'include', or variations such as 'comprises' or 30 'comprising', 'includes' or 'including' will be understood to imply the inclusion of a stated 31

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integer or group of integers, but not the exclusion 1 2 of any other integer or group of integers. 3 4 Unless the context demands otherwise, the term 5 peptide, polypeptide and protein are used 6 interchangeably to refer to amino acids in which the amino acid residues are linked by covalent peptide 7 8 bonds or alternatively (where post-translational 9 processing has removed an internal segment) by 10 covalent di-sulphide bonds, etc. The amino acid 11 chains can be of any length and comprise at least 12 two amino acids, they can include domains of 13 proteins or full-length proteins. Unless otherwise 14 stated the terms, peptide, polypeptide and protein 15 also encompass various modified forms thereof, 16 including but not limited to glycosylated forms, phosphorylated forms etc. 17 18 19 The term interaction or interacting as used herein 20 means that two entities, for example, distinct 21 peptides, domains of proteins or complete proteins, 22 exhibit sufficient physical affinity to each other 23 so as to bring the two interacting entities 24 physically close to each other. An extreme case of 25 interaction is the formation of a chemical bond that results in continual, stable proximity of the two 26 27 entities. Interactions that are based solely on 28 physical affinities, although usually more dynamic 29 than chemically bonding interactions, can be equally 30 effective at co-localising independent entities. 31 Physical affinities include, but are not limited to, 32 for example electrical charge differences,

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1	hydrophobicity, hydrogen bonds, van der Waals force,
2	ionic force, covalent linkages, and combinations
3	thereof. The interacting entities may interact
4	transiently or permanently. Interaction may be
5	reversible or irreversible. In any event it is in
6	contrast to and distinguishable from natural random
7	movement of two entities. Examples of interactions
8	include specific interactions between antigen and
9	antibody, ligand and receptor etc.
10	
11	Brief description of the figures
12	
13	The present invention will now be described with
14	reference to the following non-limiting examples and
15	with reference to the figures, wherein:
16	
17	Figure la is a ribbon diagram of EGFP;
18	
19	Figure 1b is an illustration of the split
20	points and the related sequences surrounding
21	these split points of EGFP;
22	
23	Figure 2 is a representation of a hapto-EGFP
24	with a range of linker lengths between the bait
25	peptide and respective fluorogenic fragment and
26	a plurality of peptides linked to a
27	complementary fluorogenic fragment;
28	
29	Figure 3 shows fluorescent images of Vero cells
30	transiently cotransfected with haptoEGFP
31	expression constructs, (A) Cells cotransfected
32	with pN157(6)zip and pzip(4)C158 in which a

1	functional leucine zipper mediates the
2	association of haptoEGFP1-157 and haptoEGFP158-
3	238 to generate fluorescence, (B) Negative
4	control cotransfection using $pN157(6)$ and
5	p(4)C158 which lack sequences encoding the
6	leucine zippers and as such fail to generate
7	fluorescence, (D) Cells cotransfected with
8	pN172(6) zip and $pzip(4)C173$ in which a
9	functional leucine zipper mediated association
10	of haptoEGFP1-172 and haptoEGFP173-238 occurs
11	to generate fluorescence which is of greater
12	intensity to that observed with the 157/158
13	split point (E) Negative control
14	cotransfection using $pN172(6)$ and $p(4)C173$
15	which lack sequences encoding the leucine
16	zippers and as such fail to generate
17	fluorescence, (C and F) Confocal images of
18	cotransfected cells from (A) and (D) showing
19	the intracellular localisation of fluorescence
20	- Vero cells were cotransfected with plasmids
21	encoding linkers ranging in length from $4$ to $26$
22	amino acids and UV images were collected at 24
23	hours post-transfection using identical
24	exposure times, (G) pN157( $6$ )zip and
25	pzip(4)C158 (H) $pN157(16)zip$ and $pzip(14)C158$
26	(I) $pN157(26) zip and pzip(24)C158 (J)$
27	pN157(26) zip and $pzip(4)C158$ (K) $pN157(6)$ zip
28	and pzip(24)C158 (L) a negative untransfected
29	control illustrates the background fluorescence
30	level (Italicised figures in brackets indicate
31	the length of the hydrophilic linker); and
32	

Figure 4 shows the importance of relative orientations of the haptoEGFP and binding proteins - figure 4A illustrates the case of

29

associating membrane proteins where a Type I and Type II protein combine, both hapto EGFP

6 moieties must be on the same side of the

7 membrane barrier for their combination,

8 association of membrane proteins of the same

9 type suffer from the same constraints (figure

10 4b) wherein to obtain fluorescence fusion to

the appropriate (cytoplasmic )terminus of the.

binding protein is to the same type of terminus

on both haptoEGFPs (ie: N and N' or C and C',

for Type II and Type I respectively)

15

16 Functional association of fragments of fluorescent

17 proteins, brought together by the interaction of

18 peptides fused to the fragments to screen for

19 peptide to peptide interactions requires that the

20 fragments reliably functionally associate only after

21 interaction of the fused peptides. Fluorescence may

22 be measured by suitable method known to a person

23 skilled in the art, for example, fluorescence

24 spectrometry, lunminescence spectrometry,

25 fluorescence activated cell analysis, fluorescence

26 activated cell sorting, automated microscopy or

27 automated imaging.

28

29 Reliable functional association has to date not been

30 achieved due to the possibility of steric hindrance

31 and steric constraints on the functional association

of haptoFPs when bulky proteins are associated to

30

1 the fluorescent protein fragments due to too short 2 linkers being interposed between the peptide of the 3 interest and the fragment of fluorescent protein or 4 too much flexibility due to too long a linker being 5 interposed between the same. 6 7 The inventors have determined an economical and 8 reliable method to provide a range of bait fusion 9 proteins comprising a linker region of varying length and thus provide a robust screening 10 interaction system and method. 11 12 13 This minimises the problems of steric hindrance, as 14 . a peptide of interest is provided with both 15 considerable flexibility due to the presence of long 16 linkers, but also ensures that short linkers are 17 present such that the fragments of fluorescent protein are brought into close proximity with each 18 Thus the chance of a false negative result 19 20 being obtained, i.e. finding that the peptides of study do not bind when in fact they do, is reduced. 21 22 23 A general description of the principle of the 24 invention is shown in figure 2 using haptoEGFPs as 25 the fluorescent fragments. 26 27 As shown in figure 2 protein to protein interaction 28 searches can be conducted by library interrogation. 29 The two peptides being tested for interaction are 30 designated bait and 'prey' "W". Two libraries are 31 generated (I and II), one series of constructs (here 32 designated T...Z, library I, >10,000 members) encodes

31

a hapto-EGFP followed by a DNA sequence encoding a 1 2. 60 residue linker attached to the 5'-end of a cDNA library, which contains the gene encoding the 3 4 'prey', "W" here. The second series of constructs 5 (a...e here, library II, <20 members) encodes the complementary hapto-EGFP followed by a degenerate 6 7 linker DNA sequence and the 'bait' gene. All arrows 8 indicate the direction of the polypeptide backbone (N->C). 9 10 A. 'Prey' identification: co-transfection with the 11 'prey' library (I) and construct 'e' (long linker -12 13 preferably 60 amino acid residues) from the 'bait' 14 library (II) generates fluorescent cells when the recipient cell receives a vector from library (I) 15 16 bearing the 'W' gene (in this case) and a second vector bearing the 'e' bait construct. Clonal 17 18 expansion of these fluorescent cells allows identification of gene 'W'. 19 20 21 B. Proximity measurement: The clone(s) from step A 22 are co-transfected with the 'bait' library (II). In 23 this case cells showing fluorescence synthesise interacting proteins with a sufficiently long linker 24 25 to allow productive complementary hapto-GFP interaction. ('d' or 'e' in this case), as shown to .26 the left of the diagram. The hollow arrows in the 27 right hand part of the diagram are intended to 28 indicate that the interaction of the gene products 29 with these two constructs generates fluorescence, 30 while other interactions between the product of gene 31

'W' and the bait protein do not give rise to

32

fluorescent cells due to insufficient length of 1 2 linker.

3

32

## Generation of fluorescent fragments

4 5 Fluorescent fragments may be provided by any means 6 known in the art. A first fragment of fluorescent 7 8 protein may be an N terminal fragment of fluorescent protein, e.g. GFP, comprising a substantially 9 continuous stretch of amino acids from amino acid 10 11 number 1 to amino acid X of fluorescent protein and a second fragment may be a substantially continuous 12 13 stretch of amino acids from X+1 to around the C terminal end of the fluorescent protein (e.g. amino 14 acid 238 of GFP), wherein the bond between residue X 15 and X+1 typically is located in a hydrophilic loop 16 17 region of the fluorescent protein. Should greater than two fragments of fluorescent protein require to 18 19 be generated for use in assay methods where three or 20 more fragments of fluorescent protein are linked to proteins of interest then a N terminal fragment may 21 comprise a substantially continuous stretch of amino 22 acids from amino acid number 1 to amino acid X of 23 fluorescent protein, a second fragment can be 24 25 considered as a substantially continuous stretch of amino acids from X+1 to residue Y and a third 26 fragment may be provided by a substantially 27 continuous stretch of amino acids from Y+1 to around 28 the C terminal end (e.g. amino acid 238) of 29 fluorescent protein. In such an example the bonds 30 31 between X and X+1 and Y and Y+1 will be located in

hydrophilic loop regions of fluorescent protein.

33

1 Generation of linkers 2 3 As shown in figure 2, multiple bait fusion peptides 4 may be created with linkers of differing lengths. 5 6 To enable economical extension of a linker, to 7 provide linkers of differing lengths, each linker 8 may be created using overlapping oligonucleotides 9 encoding repeating (GGGGS)x units wherein the linker oligonucleotide is engineered to carry a unique 10 restriction site, for example unique Sac I and BamHI 11 12 restriction sites, present in a core expression vector, for example pNEGFP (Sac) zip and pzip (Bam) CEGFP 13 14 (Sac I for the hexapapeptide and BamH I for the 15 tetrapeptide in example 2 ). 16 17 This allows the insertion of synthetic 18 oligonucleotides encoding further flexible 19 hydrophilic linker sequences of the form (GGGGS)<sub>n</sub> 20 with the appropriate 5' and 3' sticky ends to 21 distance the binding peptides (for example leucine 22 zippers - see example 2) away from the signalling 23 haptoEGFPs. 24 25 Once prepared the constructs may be sequenced before 26 transfection to confirm correct orientation of the 27 insert. 28 29 Further as illustrated in figure 2, a library of 30 prey fusion peptides may be provided wherein the 31 linkers of the prey fusion peptides are of the same 32 length, but different second peptides of interest

34

1 are fused to the linker region fused to the complementary fragment of fluorescent protein. 2 3 . 4 In general to prepare a library of fusion proteins 5 of unknown library sequences, the sequence encoding the hapto-EGFP is fused to the 5' end of the peptide 6 7 library due to the presence of downstream stop 8 codons in the cDNA. 9 If the gene sequence encoding the protein is 10 11 unknown, constructs are required to be generated for all three reading frames to ensure that one is in 12 13 the correct reading frame. 14 The cDNA sequences should be obtained from a source 15 which permits directional cloning into restriction 16 sites which are extremely rare in mammalian DNA. 17 Suitable sequences may be found in the Large-Insert 18 19 cDNA library (Clontech). 20 In particular embodiments a core panning vector may 21 be engineered from existing plasmids to contain a 22 CMV promoter, an initiation codon, sequences 23 24 encoding a hapto-EGFP, an intervening linker, an Sfi IA site and an Sfi IB site, a stop codon and an SV40 25 polyadenylation signal. Two additional screening 26 27 vectors may be generated to include one and two extra nucleotides between the linker and the Sfi IA 28 site to correct the reading frame. cDNA fragments, 29 30 flanked with Sfi IA and Sfi IB sites obtained from the library are cloned downstream of the optimised 31 hapto-EGFP linker constructs. The hapto-EGFP library 32

35

is then transfected into suitable cells, for example 1 CHO cells and a mixed population of cells selected 2 using G418 and passaged to confluency 3 4 Where interaction between the peptides being 5 screened occurs and the linkers allow association of the fragments of fluorescent protein, fluorescence 7 is generated. 8 9 Any cells which fluoresce may then be isolated by 10 fluorescent laser microdissection and single cell 11 RT-PCR performed to identify mRNA which encodes 12 peptides which interact with the cytoplasmic tails 13 of the receptor molecules. 14 15 Example 1 - Generation of GFP Fragments 16 17 The GFP fragments of the interaction system capable 18 of functional association were generated by split 19 points at various points along the 239 residue 20 length of the GFP protein, resulting in the 21 generation of new C' and N' termini which, in three 22 dimensions, are located at the top and at the base 23 24 of the beta-can structure. 25 Split points were introduced based on a structure 26 driven approach between hydrophilic residues. 27 28 As shown in Figure 1 the beta-can topology of EGFP 29 is formed by the eleven strands of the beta 30 structure. This structure is characterised by 31 32 forming three instances of a tripartite antiparallel

36

1 sheet motif joined edge to edge around the periphery

of the 'can', with the remaining two beta strands

completing the cylindrical structure. The most

successful split points obtained to date occur in

5 the third tripartite motif between hydrophilic

6 residues allowing adjacent hydrophobic side chains

to promote refolding of the haptoGFPs.

7 8 9

3

4

As shown in the non exhaustive list of Table 1 a

10 number of split points were identified using the

11 above approach. It would appear that each split

12 point in Table 1 is simply one example of a range of

13 potentially useful split points, the range being

shown in parentheses of Table 1.

Table 1

Split point	Residue	Possible
Number	position in	range
	EGFP	
1	23/24	(23-25)
2	38/39	(36-41)
3	50/51	(48-54)
4	76/77	(75-91)
5	89/90	(75-90)
6	102/103	(101-103)
7	116/117	(115-118)
8	132/133	(129-143)
9	142/143	(129-143)
10	157/158	(155-160)
11	172/173	(171-175)
12	190/191	(187-199)

37

13	211/212	(207-218)
14	214/215	(207-218)

1 2

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19 20

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2425

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2728

To extend the versatility of the hapto-EGFP method, constructs were created where instead of using C' and N' for the attachment of heterologus proteins, the endogenous termini, N or C, together with one of the new N' or C' termini were used (C' and N' are those N and C termini created on splitting the GFP protein into fragments, C' is thus equivalent to the new C terminal produced on the first fragment and N' is equivalent to the new N terminal produced on the complementary fragment). Using this technique the bait and prey peptides were added such that they were orientated to the associated fluorogenic fragments in the same direction as each other, for example both peptides of interest were attached to the bottom of the  $\beta$ -can structure of GFP or in the opposite direction, for example the bait peptide was attached to the bottom of the  $\beta$ -can structure of GFP, while the prey protein was attached to the top of the  $\beta$ -can structure of GFP. As shown in figures 4 A & B, as peptides interact with each other in a particular orientation, then the direction of the linkage of the peptide to the N, N', C or C' end of the fluorogenic fragment may be important in certain circumstances so as to allow the fluorescent protein fragments to functionally interact following interaction of the peptides.

29

1	Example 2
2	
3	To determine the effect of varying the length of the
4	intervening hydrophilic linkers interposed between
5	complementary fragments of fluorescent protein and
6	leucine zipper proteins known to bind to each other
7	the linkers were empirically increased in length in
8	decapeptide units using the general method detailed
9	above to modify linkers of both pN157(6)zip and
10	pzip(4)C <sup>158</sup> to increase the linker by 10, 20, 30 and
11	40 residues by the insertion of complementary
12	oligonucleotides with Sac I and BamH I sites to
1.3	generate in the case of the $N^{157}(6)$ zip chimera, to
14	16, 26, 36 and 46 and, in the case of the
15	complementary zip(4)C158 chimera, to 14, 24, 34 and
16	44 residues.
17	
18	The results of this study are shown in figure 3.
19	
20	No significant differences in the levels of
21	fluorescence were observed when the hydrophilic
22	spacers were lengthened by up to 26 and 24 amino
23	acids respectively. However, the signal increased
24	when spacers of 36 and 34 separated the leucine
25	zipper and the haptoEGFP moieties, whereas the
26	signal decreased when linkers comprised of 46 and 44
27	amino acids were introduced.
28	
29	Example 3
30	
31	Utilisation of MV H as a model homo-oligomerising
32	transmembrane glycoprotein

39

1 2 In order to demonstrate that this approach can be 3 used for a wider range of applications than current reporter systems the membrane glycoproteins of Measles Virus (MV) were examined. 5 6 7 Measles virus (MV) infection is mediated by a complex of two viral envelope proteins, 8 9 haemagglutinin (H) glycoprotein and fusion (F) glycoprotein that bind to each other and then 10 11 complex with surface receptors to aid the fusion of the virus with the plasma membrane. 12 glycoprotein is dimerised in the endoplasmic 13 14 reticulum and is thought to exist on the cell 15 surface as a tetramer (dimer of dimers). The fusion (F) glycoprotein, is synthesised as an inactive 16 17 precursor (F<sub>0</sub>) which is a highly conserved type I 18 transmembrane glycoprotein of about 60kDa, which is cleaved by furin in the trans-golgi to yield the 19 41kDa (f<sub>1</sub>) and the 18kDa (f<sub>2</sub>) disulphide-linked 20 21 activated F-protein. Infection of the measles virus 22 is dependent on the interaction of the F/H complex 23 with cell surface receptors. 24 25 Two constructs, which expressed N157(16)MV-H and C158(14)MV-H, were initially generated in order to 26 27 investigate homodimerisation of a type II membrane 28 glycoprotein of unknown structure. The linker 29 regions of these constructs were generated using 30 overlapping oligonucleotides which contain Sfi IA and Sfi IB restriction sites were introduced into 31  $pN^{1/157}(16)$  zip and  $pC^{158/239}(14)$  zip constructs. These 32

1 chimeras differ from those generated from the

2 leucine zippers in that the first has H fused to the

40

3 C' terminus, while the second employs the endogenous

4 C terminus for fusion. Expression of the chimeric

5 proteins was detected by immunoblotting cell lysates

6 using peptide antiserum raised against EGFP (results

7 not shown). This demonstrated that the haptoEGFP

8 tagged H glycoproteins were stably expressed in the

9 transfected cells. Furthermore, the electrophoretic

10 mobility of the chimeric proteins suggested that

11 they were correctly glycosylated. Fluorescence was

12 readily detected in living cells and all of the

13 necessary controls demonstrated that the association

of the haptoEGFPs was specifically driven by the

dimerisation of the H glycoproteins. Fluorescence

16 was absent from the nucleus but was clearly

17 demonstrable from the ER through the Golgi to the

18 plasma membrane of the cells.

19

20 It is clear that this methodology could be used to

21 identify further, membrane receptor proteins which

interact with the H protein as could cytoplasmic

23 proteins which interact with known MV receptors and

24 which may therefore initiate downstream signalling

25 events.

26 27

Example 4

28

29 In order to ascertain that the haptoEGFP tagged

30 glycoproteins were capable of forming a biologically

31 active complex at the cell membrane cells were

32 transfected with constructs expressing a number of

1

41

different H and F chimeras. Firstly the bioactivity

of the H chimeras was investigated by co-2 3 transfection with a plasmid expressing the unmodified F glycoprotein. Initially cell-to-cell 4 fusion was readily detected 2 d.p.t. in cells 5 6 expressing N157(16)MV-H, C158(14)MV-H, and F. 7 8 Multi-nucleated syncytia comprised of more that 50 9 cells were obtained which were easily discernible by phase-contrast microscopy. 10 11 12 Fluorescence was detected by vital confocal laser microscopy in all syncytia, their size was 13 14 comparable to that obtained by co-expression of 15 unmodified MV F and H. 16 By three days post-transfection, cell-to-cell fusion 17 18 was detected over large areas of the monolayer and many syncytia comprised of over 200 individual 19 cells. Confocal scanning laser microscopy was used 20 21 to determine whether localised fluorescence was present within the syncytia and series of images 22 23 were collected. Composite images were generated and 24 fluorescence localization was examined in the x/z 25 and y/z planes. Fluorescence was detected in the perinuclear regions and also in a honeycomb lattice 26 27 which is consistent with the presence of the glycoprotein in the ER and Golgi. 28 29 30 When the plasma membrane was examined in x/z and y/z31 it was difficult detect a discrete line of 32 fluorescence in single sections. However, small 1-5

42

1 µm vesicles with fluorescent membranes were 2 frequently detected at the cell surface. These vesicles are very reminiscent of budding virions and 3 are approximately 10 times larger than MV virions. 5 6 These co-transfected cells were fixed in order to 7 examine the intracellular localisation of 8 fluorescence within syncytia at higher 9 magnifications. In the fixed cells it was also 10 clear that the fluorescence was present in the ER 11 and Golgi as expected. However, areas of localised 12 fluorescence were also detected at the periphery of the syncytia where the fused cells came into contact 13 14 with the surrounding cells, suggesting that the H 15 glycoprotein dimers are not evenly distributed on 16 the plasma membrane and these accumulations could be 17 sites of fusion pore formation where the H 18 glycoproteins are in close contact with the cellular 19 receptor, in this case CD46. 20 21 The extracellular localisation of the H dimers was 22 also examined by indirect immunofluorescence using 23 an anti-H MAb on unpermeabilised cells. vital immunostaining indicated that a significant 24 25 percentage of the dimeric H chimera had been 26 correctly processed and trafficked to the cell membrane where, in view of the size of the syncytia, 27 28 it was clearly functional. Fluorimetery was used to 29 determine if the fluorescence could be detected and 30 quantified. In cells transfected for defined periods of time it was found that syncytia formed. 31 Fluorescent signals were detected which were 32

Τ.	equivalent to those obtained in phis/(6/21p and
2	pzip(4)C158 co-transfected cells. No signals were
3	obtained when the construct which expressed
4	C158(14)MV-H was replaced by one encoding
5	zip(14)C158 indicating that the specific association
6	of the H glycoproteins was driving the haptoEGFP
7	moieties into close enough proximity to enable the
8	generation of the fluorophore.
9	
10	Although the invention has been particularly shown
11	and described with reference to particular examples,
12	it will be understood by those skilled in the art
13	that various changes in the form and details may be
14	made therein without departing from the scope of the
15 <sup>.</sup>	present invention.
16	
17	
18	